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54 Apparatus for the simultaneous synthesis of different peptides.

57 The present invention is directed to the simultaneous multiple chemical synthesizer comprising a number of reaction vessels wherein each vessel has a filter in the bottom portion thereof, a number of needles wherein each needle is connected to a pair of an aspiration injection line of a reaction mixture and a gas supply line in charge of each reaction vessel and each needle does not touch resin, a number of arms which are horizontally and vertically movable and hold the respective needles, a pair of a bubbling gas line and a waste discharge line in charge of each reaction vessel, wherein each line is connected to the bottom portion of each reaction vessel, a number of purge portions which move synchronously with said waste discharge lines, and means for washing the contact portions with the reaction reagents of the needles and the purge portions.

EP 0 529 504 A2

## FIELD OF THE INVENTION

The present invention relates to a simultaneous multiple chemical synthesizer.

## 5 BACKGROUND OF THE INVENTION

In conventional solid-phase synthesizers, particularly multiple synthesizers wherein a single needle is in charge of all of a number of reaction vessels, repeated reaction tends to cause incomplete reaction, cross contamination more times in moving needle and the like. This problem arises due to the different reactivities  
 10 of individual chemical derivatives in the case of multiple reactions. In addition, from the viewpoint of efficiency-related factors, such as yield and purity of reaction product, it is disadvantageous to react all items by the same method or under the same conditions. Moreover, the single needle must be frequently moved reciprocally among a large number of reaction vessels, various reagent containers and washing ports, which results in increased operating time or side reactions. Specifically, when a single needle is in  
 15 charge of a number of reaction vessels, while the needle is acting on one reaction vessel, the other reaction vessels must wait, and in addition, washing must be completed by the reach of the needle to another reaction vessel, which results in increased time requirement for each reaction. This method is also disadvantageous from the viewpoint of the mechanical strength and durability of the apparatus because it involves a large number of movements. Moreover, when a needle contacts with resin by the bubbling or the  
 20 stirring, the resin attaches to the needle, thereby causing the loss of resin and cross contamination.

In the apparatuses wherein the reagents in respective reaction vessels are aspirated and discharged on a one-by-one basis by a needle, not only the discharge of the reagents, washing solvents, etc. is incomplete, but also thorough washing is essential for each discharge, which results in cross contamination.

In the apparatuses wherein the reagents are simultaneously discharged from all reaction vessels under  
 25 reduced pressure with the bottom portion thereof kept at a negative pressure, there are some problems of difficulty in the maintenance of tightness between the vessels and the bottom portion containers and difficulty in uniform discharge due to the differences in the properties of reaction products among the items.

Also, in conventional methods, it must be outside the reaction apparatus to take out the reaction product from the solid phase by another deprotecting reaction, namely cleavage, after reaction, i.e., they are not  
 30 efficient solid-phase synthesizing methods.

## SUMMARY OF THE INVENTION

It is an object of the present invention to provide a simultaneous multiple chemical synthesizer which  
 35 allows simultaneous multiple chemical reactions with high efficiency in terms of reaction yield, purity of synthesized product, rapidness, low cost, easy maintenance, etc..

In view of the problems described above, the present inventors have made investigations and have found that it is possible to realize a high-efficiency production by using separate systems for respective  
 40 reactions, thereby solving the problems such as the reduction of reaction efficiency in conventional reaction apparatuses.

Accordingly, the present invention is directed to a simultaneous multiple chemical synthesizer, each reaction lane being arranged independently, comprising a number of reaction vessels wherein each vessel has a filter in the bottom portion thereof, a number of needles wherein each needle is connected to a pair of  
 45 an aspiration injection line of a reaction mixture and a gas supply line in charge of each reaction vessel, a number of arms which are horizontally and vertically movable and hold the respective needles, a pair of a bubbling gas line and a waste discharge line in charge of each reaction vessel, wherein each line is connected to the bottom portion of each reaction vessel, a number of purge portions which move synchronously with said waste discharge lines, and means for washing the contact portions with the reaction reagents of the needles and the purge portions.

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## BRIEF DESCRIPTION OF THE DRAWINGS

The present invention will become more fully understood from the detailed description given herein-  
 below and the accompanying drawing which is given by way of illustration only, and thus, is not limitative of  
 55 the present invention, and wherein:

Figure 1 is a schematic diagram of the simultaneous multiple chemical synthesizer of the present invention;

Figure 2 is a schematic diagram of needle and purge portions in the simultaneous multiple chemical synthesizer of the present invention;

Figure 3 is a schematic diagram of a solid-phase synthesizer with the simultaneous multiple needle of the present invention;

5 Figure 4 is a chromatogram of crude cleaved leucine enkephaline in reversed-phase HPLC which was simultaneously and independently synthesized in 8 reaction channels in Example 1, using the apparatus of the present invention;

Figure 5 is a chromatogram of HPLC of tripeptide synthesized in Example 2 using the apparatus of the present invention; and

10 Figure 6 is a chromatogram in reversed-phase HPLC of 8 different crude peptides obtained by simultaneous cleavage in Example 3 using the apparatus of the present invention.

The reference numbers in Figures 1 through 6 denote the following elements:

Element 1 is a needle, element 2 an arm, element 2' an arm, element 3 a motor, element 4 a motor, element 5 a reaction vessel, element 6 an O ring, element 7 a purge nozzle, element 8 a waste discharge  
15 line, element 9 a gas line for bubbling, element 10a a gas supply line, element 10b a gas supply line, element 11a washing port, element 12 reactant station, element 13 a bottle for washing solvent, element 14 a microsyringe for aspiration and injection and element 15 a bottle for waste liquid.

## DETAILED DESCRIPTION OF THE INVENTION

20 Figure 1 is a schematic diagram of the simultaneous multiple chemical synthesizer of the present invention.

Motors 3 and 4 are provided so that arm 2, which holds needle 1 etc., can be moved horizontally and vertically, respectively. Needle 1 is connected with a series of aspiration injection line with microsyringe 14  
25 for aspirating and injecting solvent or reagents, etc. from reactant station 12 to reaction vessel 5 and connected with gas supply line 10a for injecting the gas into reaction vessel 5. The purge portion is provided to seal the upper face of reaction vessel 5 and supply the unreacted reagents, solvents and washing solvents to waste discharge line 8 by a gas pressure exerted by gas introduction after completion of reaction or resin washing, and it is configured with purge nozzle 7, which is equipped with O-ring 6 for  
30 tight sealing. As illustrated in Figure 2, needle 1, and purge nozzle 7, which is equipped with O-ring 6 for tight sealing, are fixed to arm 2; a number of pairs thereof each of which is in charge of each reaction vessel are equipped, as illustrated in Figure 1.

In Figures 1 and 3, five reaction lanes are shown, but this number of lanes can be increased or decreased. Also, in Figures 1 and 2, needle 1 and purge nozzle 7 are attached to the same arm 2, but they  
35 may be attached to separate driving arms 2 and 2', as illustrated in Figure 3.

Desirably, reaction vessel 5 is a disposable reaction vessel having a filter in the bottom portion thereof. As an example, a polypropylene reaction vessel packed with a polyalkylene filter in the bottom portion thereof may be mentioned. Examples of polyalkylene filter materials include polypropylene, polyethylene and polyvinylidene fluoride, with preference given to polypropylene. There is no limitation on the pore size  
40 of filter or the thickness of the filter layer packed in the reaction vessel, as long as they are in the ordinary range; for example, the pore size of filter is such that the resin, which is 200 to 400 mesh (63 to 125  $\mu\text{m}$ ) in size, cannot pass the filter, and it is usually about 5 to 10  $\mu\text{m}$ .

Any material can be used for the main body of the reaction vessel, as long as it is not likely to generate static electricity on the solid-phase resin support; polypropylene is preferred, since it is cheap and shows  
45 little unspecific adsorption of the resulting peptide. Such a filter in the reaction vessel need not be pre-treated before using the reaction vessel because it is free of the problem of easy clogging. Therefore, polypropylene is suitable for simultaneous multiple chemical synthesis as in the present invention.

Figure 3 is a schematic diagram of a solid-phase synthesizer based on the simultaneous multiple chemical synthesizer illustrated in Figure 1. Reaction vessel 5 is a disposable reaction vessel having a filter  
50 in the bottom portion thereof wherein a solid-phase resin support is packed, as described above; for each reaction vessel, a pair of needle 1 and purge nozzle 7 at purge portion is driven. Any gas can be used, as long as it does not affect the reaction; for example, nitrogen gas, which is cheap, is preferred. Needles do not touch with the resin support.

Bubbling gas line 9 is for gas introduction to reaction vessel 5 via the bottom portion thereof for stirring  
55 the reaction mixture in order to accelerate and uniformize the reaction, while waste discharge line 8 is for discharging wastes such as the unreacted reagents, solvents and by-products from reaction vessel 5 via the bottom portion thereof after completion of the reaction. Bubbling gas line 9 and waste discharge line 8 are connected to the bottom portion of reaction vessel 5 via a three-way valve or another means.

Also, gas supply line 10b for discharging the reagents and solvents from the reaction vessel is connected to the above-mentioned purge nozzle 7, via which a gas is introduced to discharge the waste from the reaction vessel to bottle for waste liquid 15. A pair of bubbling gas line 9 and waste discharge line 8 exists in charge of each reaction vessel 5.

Also, an aspiration injection line with microsyringe 14, which is for aspirating the reaction mixture of solvent, reagents, etc. from reactant station 12 and injecting them into the reaction vessel, also serves as a line for injecting the washing solvent from bottle for washing solvent 13 into the reaction vessel. The microsyringe is equipped with a pair of a syringe and a motor which independently drives a piston for each reaction vessel so that different volumes of reaction mixture can be supplied to the reaction vessel of each line. Gas supply line 10a is used to cause bubbling to thoroughly mixing the starting material, reagents, solvents, catalysts, etc. set on reactant station 12 to prevent the degrading after they are injected via aspiration injection line with microsyringe 14. Said aspiration injection line with microsyringe 14 and gas supply line 10a are connected to a needle through a line via a three-way valve.

As means for washing the contact portions with the reaction reagents of needle 1 and purge nozzle 7 at the purge portion, washing port 11 is provided, where said contact portions are washed after injection of the reagent and after completion of the reaction.

Reactant station 12, bottle for washing solvent 13, etc. permit optional selection of the number thereof according to the route of synthesis. They may also be automated by microprocessor control.

As an example of the use of the simultaneous multiple chemical synthesizer of the present invention, peptide synthesis using BOP/HOBt is described below. First, resin support containing reaction vessel 5 is set in position, and HOBt (N-hydroxybenzotriazole), NMM (N-methylmorpholine), piperidine, methanol, t-butyl methyl ether and each  $N_\alpha$  protected amino acid (powder) together with PyBOP® (benzotriazole-1-yloxy-tris-pyrrolidino-phosphonium hexafluorophosphate; produced by Novabiochem A.G) are placed on reactant station 12 corresponding to the reaction vessel. Bottle for washing solvent 13 contains DMF (dimethylformamide) as a washing of a reactant dissolving solvent. The following operations are controlled by a microprocessor.

#### 1. DMF wash

DMF is injected to reaction vessel 5 via an aspiration injection line; a gas is supplied via bubbling gas line 9 to wash the resin; the upper face of the reaction vessel is closed tight at the purge portion; the DMF in reaction vessel 5 is purged into bottle for waste liquid 15.

#### 2. Piperidine wash

Needle 1 is moved to the piperidine container in reactant station 12 by means of robot arm 2, and the piperidine is aspirated into aspiration injection line, after which the needle is moved to reaction vessel 5, and the piperidine solution is injected, followed by bubbling in the same manner as in DMF wash to deprotect the  $N_\alpha$  protecting group of the first amino acid bounded on the resin. After completion of the reaction, purging is performed.

#### 3. Piperidine wash

The same as piperidine wash step 2.

#### 4. DMF wash

The same as DMF wash step 1.

#### 5. Amino acid activation

To the container which contains the amino acid together with PyBOP, BOP or TBTU (2-(1H-benzotriazole-1-yl)-1,1,3,3-tetramethyluronium tetrafluoroborate) to be next incorporation, which is set up on the reactant station, HOBt solution and NMM solution are injected by means of robot arm 2 and needle 1. After discharging the liquids from needle 1 and microsyringe 14 at washing port 11, the content in the container containing the amino acid is dissolved by bubbling with the gas supplied from the needle tip.

## 6. Coupling reaction

The amino acid activated in step 5 is transferred to reaction vessel 5 via an aspiration injection line with microsyringe 14 and bubbled via bubbling gas line 9; after completion of the reaction, purging is performed.

5

## 7. DMF wash

The same as DMF wash step 1.

The above steps are repeated in the number of cycles equivalent to the number of amino acids to be synthesized. Then, after the above step 4 is completed as the final cycle, the resin is washed with methanol and t-butyl methyl ether in the same manner as piperidine treatment, and the resin is finally dried with the nitrogen gas introduced from the purge portion.

The simultaneous multiple chemical synthesizer of the present invention is particularly suitable for the synthesis of biochemical substances such as peptides, nucleic acids and sugar chains. The apparatus of the present invention is characterized in that 1) time requirement per reaction can be shortened because the aspiration injection lines of reaction mixture are independently in charge of respective reaction vessels, because the waste discharge lines are also independent, and because each needle is in charge of each reaction vessel, 2) the problems of side reactions and cross contamination can be solved because means for washing the contact portions is provided, and 3) cleavage namely a reaction for taking out the desired reaction product from the solid phase can be carried out simultaneously in the apparatus of the present invention because the reaction vessels used have a particular filter. Also, individual reactions can be monitored because the waste discharge lines are independently arranged for respective reaction vessels.

The use of the apparatus of the present invention allows simultaneous multiple chemical reactions (e.g., solid-phase peptide synthesis) with high efficiency in terms of reaction yield, time requirement for synthesis and synthesized product purity. Also, the ease of peptide synthesis varies widely depending on the type and sequence of the constituent amino acids of the desired peptide; the use of the apparatus of the present invention allows simultaneous multiple synthesis even when different reagents and different reaction methods are used for respective reaction systems. For these reasons, in synthesizing the same desired peptide by different methods, the apparatus of the present invention makes it possible to 1) evaluate the reagents for synthesis and 2) evaluate the methods for synthesis at the same time. Furthermore, because the apparatus of the present invention is capable of simultaneously synthesizing a several types of similar compounds and related compounds, its applicability is wide and it is useful in epitope mapping, structure-activity correlation studies and screening of more active analogues.

## 35 EXAMPLES

## Example 1

Synthesis of the opioid peptide leucine enkephalin comprising 5 amino acids:

40

H-Tyr-Gly-Gly-Phe-Leu-OH was synthesized simultaneously in 8 reaction channels A through H, and the differences among the reaction channels were investigated.

Using Fmoc-Leu-p-benzyloxybenzyl alcohol resin (30 mg) as the solid support, N<sub>α</sub>-Fmoc amino acids activated with PyBOP®-HOBt were sequentially incorporated from the C-terminus in the presence of N-methylmorpholine (NMM). Before the introducing (coupling) reaction, the N<sub>α</sub>-Fmoc group was removed with 30% piperidine in DMF, followed by washing with DMF alone. Because the resin substitution rate was 0.38 meq/g, the reaction capacity of the 30 mg starting resin was 11.4 μmol. Thus, calculated yield of the desired product (molecular weight 555.64) is 6.33 mg, about 6.0 to 7 mg of cleaved peptide was obtained in each lane.

The synthesis schedule used in this reaction is shown below. The reaction apparatus, under microprocessor control, was automatically operated in this order of reaction steps for each component.

55

## [Calculation Table for Synthesis]

5

Lane : A. B. C. D. E. F. G. H

Strategy : Fmoc/Mod.BOP-HOBT

10

Peptide : Leu-Enkephalin

Sequence : Tyr-Gly-Gly-Phe-Leu-OH

15

: C<sub>28</sub>H<sub>37</sub>N<sub>5</sub>O<sub>7</sub> (mwt: 555.64)

Resin : Fmoc-Leu-p-benzyloxybenzyl alcohol resin

20

Resin Substitution : 0.38 meq/g

Resin Quantity : 30.00 mg

25

Free Peptide (theor. yield) : 6.33 mg

30

Amino Acid Station(Excess 10 fold): 114  $\mu$ mol

4:Fmoc-Phe-OH mwt= 387.44 44.17 mg

35

3:Fmoc-Gly-OH mwt= 297.31 33.89 mg

2:Fmoc-Gly-OH mwt= 297.31 33.89 mg

1:Fmoc-Tyr(tBu)-OH mwt= 459.54 52.39 mg

40

PyBOP mwt: 520.3 (  $\times$  1.0 eq) 59.31 mg /Coupling

45

NMM 110 ml/mol (  $\times$  1.5 eq) 171.0  $\mu$ l /Coupling

1.0 mmol/ml in DMF

HOBT mwt: 135.1 (  $\times$  1.0 eq) 228.0  $\mu$ l /Coupling

50

0.5 mmol/ml in DMF

55

## [Synthesis Schedule]

	[Step]	[Operation]	[Times]
5			
	1	DMF Wash	1 min. × 1
10	2	Piperidine (30 % in DMF) Wash	5 min. × 1
	3	Piperidine (30 % in DMF) Wash	3 min. × 1
	4	DMF Wash	1 min. × 5
15	5	Activation of Amino Acids	1 to 3 min
	6	Coupling(Mixing with Resin)	30 min.
20	7	DMF Wash	1 min. × 4

## [Final Cycle]

	[Step]	[Operation]	[Times]
25			
	1 0 0	DMF Wash	1 min. × 1
30	1 0 1	Piperidine Wash	5 min. × 1
	1 0 2	Piperidine Wash	3 min. × 1
	1 0 3	DMF Wash	1 min. × 5
35	1 0 4	Methanol Wash	1 min. × 2
	1 0 5	t-butylmethylether	0.5min. × 1
40	1 0 6	Nitrogen Blow	10 min.

In this case, 3 bottles were prepared; piperidine (30 % in DMF), methanol and t-butylmethylether. On the reactant station, each amino acid was weighed with PyBOP® as shown in calculation table, Fmoc-Phe-OH, Fmoc-Gly-OH, Fmoc-Gly-OH, Fmoc-Tyr(tBu)-OH, NMM solution in DMF and HOBt solution in DMF were placed; the gas used was gaseous nitrogen. The reaction vessel used was of polypropylene, packed with a polypropylene filter therein.

After completion of the peptide chain elongation, the resulted protected peptide resin having the N<sub>α</sub>-Fmoc group was treated with 30 % piperidine in DMF and then was cleaved (cleavage; reaction simultaneously removal of side chain protecting groups with peptide chain removal from support) with trifluoroacetic acid (TFA) containing 5% anisole and 1% ethanedithiol while remaining in the reaction vessel; the resulting mixture was precipitated with diethyl ether and centrifuged. The resulting precipitate was dissolved in 10% acetic acid and lyophilized to yield a crude pentapeptide. The following shows common procedures of cleavage after synthesis and of free peptide recovery. This reaction was performed in each reaction vessel used for the synthesizing reaction at the same time. Accordingly,

Dry the reaction vessel under reduced pressure →

Remove the protective group (cleavage cocktail 0.3 to 0.5 ml/reaction vessel)→

Mix slowly →

Filter while blowing nitrogen gas from the upper portion of the reaction vessel →

Collect the filtrate (0.3 to 0.5ml) in centrifugal tube →

Precipitate with cold diethyl ether (15 ml) →

Centrifuge (3000 rpm, 5 minutes) →

5 Decanted, wash with diethyl ether and centrifuge twice →

Dry with nitrogen gas stream for 1 to 2 minutes →

Dissolve in 30% acetic acid (0.5 ml) →

Dilute with water (2 to 3 ml) →

Lyophilize →

10 Obtain a crude peptide .

Yield was constant at  $6.5 \pm 0.5$  mg among the 8 channels. The results of reversed-phase HPLC are shown in Figure 4. The analytical conditions for the reversed-phase HPLC were as follows.

Column : SynProPep<sup>®</sup> RPC18 (4.6×250 mm) (\* = Trademark)

Eluent : 0.01N HCl /CH<sub>3</sub>CN = 85/15 to 55/45 (30 min.)

15 Flow Rate : 1.0 ml/min. Absorbance : 210 nm

There was no difference among channels A through H, and high purity was shown. The peptide obtained was identified as the desired product by sequence analysis and mass spectrometry.

## Example 2

20

Synthesis by the simultaneous different coupling method:

The tripeptide H-Gly-His-Lys-OH, a hepatocyte growth factor, was synthesized simultaneously by the following 4 coupling methods A through D. The starting resin used was Fmoc-Lys(Boc)-p-benzyloxybenzyl

25 alcohol resin.

A: Method using DICD (diisopropylcarbodiimide), the acyl component converted to an active ester by means of HOBt was reacted with the amino component on the solid-phase resin support.

B: Method using PyBOP<sup>®</sup> and HOBt (the same as in Example 1 described above).

30 C: Method using the uronium salt TBTU (2-(1H-benzotriazole-1-yl)-1,1,3,3-tetramethyluronium tetrafluoroborate) in place of PyBOP<sup>®</sup>.

D: Method using Pfp Ester (pentafluoro ester of amino acid).

In any case, synthesis was performed on the schedule shown in Example 1, and cleavage was performed using the reaction vessel as such by the method described in Example 1. The HPLC pattern of tripeptide obtained are shown in Figure 5. The analytical conditions for the reversed-phase HPLC were as

35 follows.

Column : SynProPep<sup>®</sup> Pep Kat (4.0×50 mm)(\* = Trademark)

Eluent : 5mM NaH<sub>2</sub> PO<sub>4</sub> /5mM NaH<sub>2</sub> PO<sub>4</sub> + 1M NaCl = 100/0 to 75/25 (30 min.)

Flow Rate : 1.0 ml/min. Absorbance : 210 nm

The resulted peptide also shows good efficiency and satisfied by sequence analysis, etc..

40

## Example 3

Simultaneous multiple synthesis:

45 Using 8 independent channels, peptides with respectively different amino acid sequences were simultaneously synthesized and cleaved to provide 8 crude desired peptides with yields close to the theoretically calculated values respectively. These synthesis were achieved in accordance with the schedule and cleavage procedure as described in Example 1.

50 In this synthesis, the cleavage cocktail used were: 94% TFA (trifluoroacetic acid), 5% anisole and 1% ethanediol (EDT) (90 min); 94% TFA, 3% anisole, 3% EDT and 5mg of 2-methylindole (90 min) for Trp containing peptides; 82% TFA, 5% water, 5% thioanisole, 3% EDT 2% methylethylsulfide and 3% phenol (6 hrs) for Arg containing peptides.

The 8 different crude peptides (lanes A through H), the method for coupling the starting resin and other data are shown below.

55 Fmoc Amino Acid : 10 fold excess against amino component Coupling PyBOP<sup>®</sup> -HOBt-NMM (1:1:1.5)

The side chain protection groups used were :

Asp = OBut; Arg = Pmc; Asn, His = Trt; Ser, Thr, Tyr = tBu



Starting resin : PAL™ resin (50 mg, the substitution group = 0.21 mmol/g, 10.5 μmol)

Lane A : Rat Neuromedin (16-25) (MW = 1208.42) H-Gly-Gly-Phe-Phe-Leu-Phe-Arg-Pro-Arg-Asn-NH<sub>2</sub>

Lane B : Porcine S.C.Neuromedin-K (MW = 1209.44) H-Asp-Met-His-Asp-Phe-Phe-Val-Gly-Leu-Met-NH<sub>2</sub>

5 Lane C : Porcine S.C.Neuromedin-L (MW = 1132.34) H-His-Lys-Thr-Asp-Ser-Phe-Val-Gly-Leu-Met-NH<sub>2</sub>

Lane D : Porcine S.C.Neuromedin-B (MW = 1131.31) H-Gly-Asn-Leu-Trp-Ala-Thr-Gly-His-Phe-Met-NH<sub>2</sub>

10 Lane E : Porcine S.C.Neuromedin-C (MW = 1119.30) H-Gly-Asn-His-Trp-Ala-Val-Gly-His-Leu-Met-NH<sub>2</sub>

Lane F : Porcine S.C.Neuromedin-N (MW = 743.98) H-Lys-Ile-Pro-Tyr-Ile-Leu-NH<sub>2</sub>

Lane G : Porcine S.C.Neuromedin-U8 (MW = 1110.32) H-Tyr-Phe-Leu-Phe-Arg-Pro-Arg-Asn-NH<sub>2</sub>

Lane H : Porcine S.C.GRP-10 (MW = 1119.30) H-Gly-Asn-His-Trp-Ala-Val-Gly-His-Leu-Met-NH<sub>2</sub>

15 For peptide synthesis on lane A, for instance, a 30% piperidine solution in DMF as an N<sub>α</sub>-deprotecting reagent, DMF, methanol, t-butyl methyl ether, etc. as bottle for washing solvents, and an NMM solution in DMF and an HOBt solution in DMF for activation reaction were prepared. On the reactant station, mixtures of each of various protected amino acid derivatives together of PyBOP® such as Fmoc-Gly-OH, Fmoc-Gly-OH, Fmoc-Phe-OH, Fmoc-Phe-OH, Fmoc-Leu-OH, Fmoc-Leu-OH, Fmoc-Phe-OH, Fmoc-Arg(Pmc)-OH, Fmoc-Pro-OH, Fmoc-Arg(Pmc)-OH and Fmoc-Asn(Trt)-OH were arranged in the order of coupling. For the other lanes as

20 well, these materials were arranged in the order of coupling. The gas used was gaseous nitrogen. In cases where a peptide with less amino acids than on other lanes that is shorter fragment is synthesized as on lanes F and G, no vials containing amino acid derivatives were placed and no reagents were supplied in accordance with the program of synthesis.

25 The patterns of reverse-phase HPLC of 8 different crude peptides obtained by simultaneous cleavage using cleavage cocktail described above are shown in Figure 6. The analytical conditions for the reversed-phase HPLC were as follows.

Column : SynProPep® RPC18 (4.6×250 mm) (\* = Trademark)

Eluent : 0.01N HCl /CH<sub>3</sub>CN = 85/15~ 55/45 (30 min.)

Flow Rate : 1.0 ml/min., Absorbance at 210 nm

30 The results of FAB mass spectra at a main peak were also described here. All results show high purity and the results of mass spectra were calculated value.

Moreover, the peptides obtained here were confirmed their primary structure by sequence analysis.

The peptides obtained here, namely obtained by simultaneous multiple synthesis, also show high homogeneity by sequence analysis.

35 The present invention being thus described, it will be obvious that the same may be varied in many ways. Such variations are not to be regarded as a departure from the spirit and scope of the invention, and all such modifications as would be obvious to one skilled in the art are intended to be included within the scope of the following claim.

#### 40 Claims

1. A simultaneous multiple chemical synthesizer, each reaction lane being arranged independently, comprising:

a number of reaction vessels wherein each vessel has a filter in the bottom portion thereof;

45 a number of needles wherein each needle is connected to a pair of an aspiration injection line of a reaction mixture and a gas supply line in charge of each reaction vessel, and each needle does not touch resin in the reaction vessel;

a number of arms which are horizontally and vertically movable and hold the respective needles;

50 a pair of a bubbling gas line and a waste discharge line in charge of each reaction vessel, wherein each line is connected to the bottom portion of each reaction vessel;

a number of purge portions which move synchronously with said waste discharge lines; and

means for washing the contact portions with the reaction reagents of the needles and the purge portions.

55

\* 5-(4'-(9-Fluorenylmethyloxy-carbonyl) aminomethyl-3,5-dimethoxyphenoxy)-valeric  
acid handle attached to p-methylbenzhydryl-amine resin

2. The simultaneous multiple chemical synthesizer according to claim 1, further comprising means for introducing gas to each reaction vessel so as to discharge the waste from the bottom portion of each reaction vessel to a bottle for waste liquid.
- 5 3. The simultaneous multiple chemical synthesizer according to claim 1, wherein said reaction vessel has a polyalkylene filter selected from the group consisting of polypropylene, polyethylene and polyvinylidene fluoride.
- 10 4. The simultaneous multiple chemical synthesizer according to claim 1, wherein a pore size of the filter is about 5 to 10 $\mu$ m.

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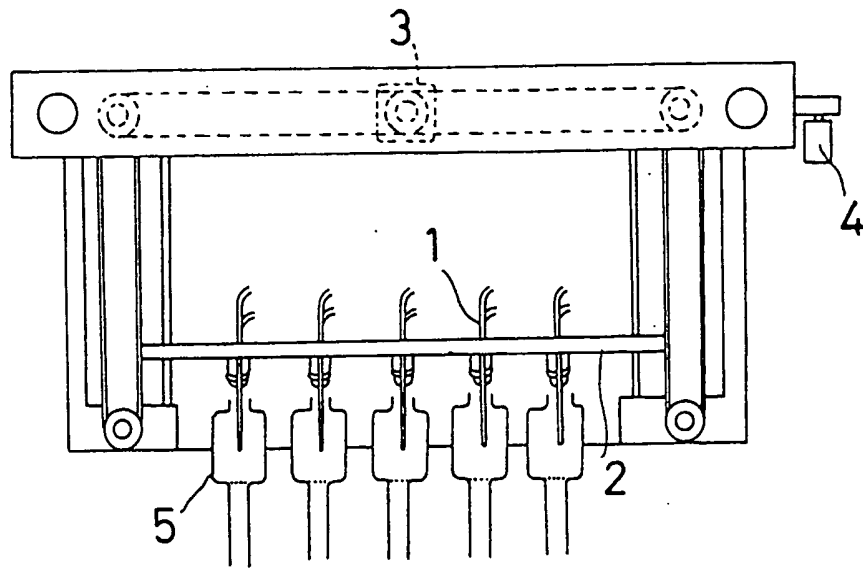
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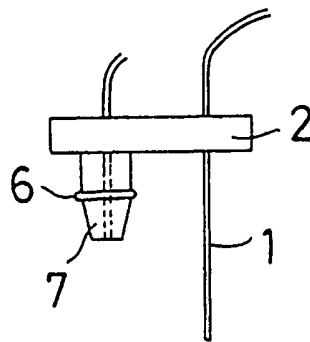
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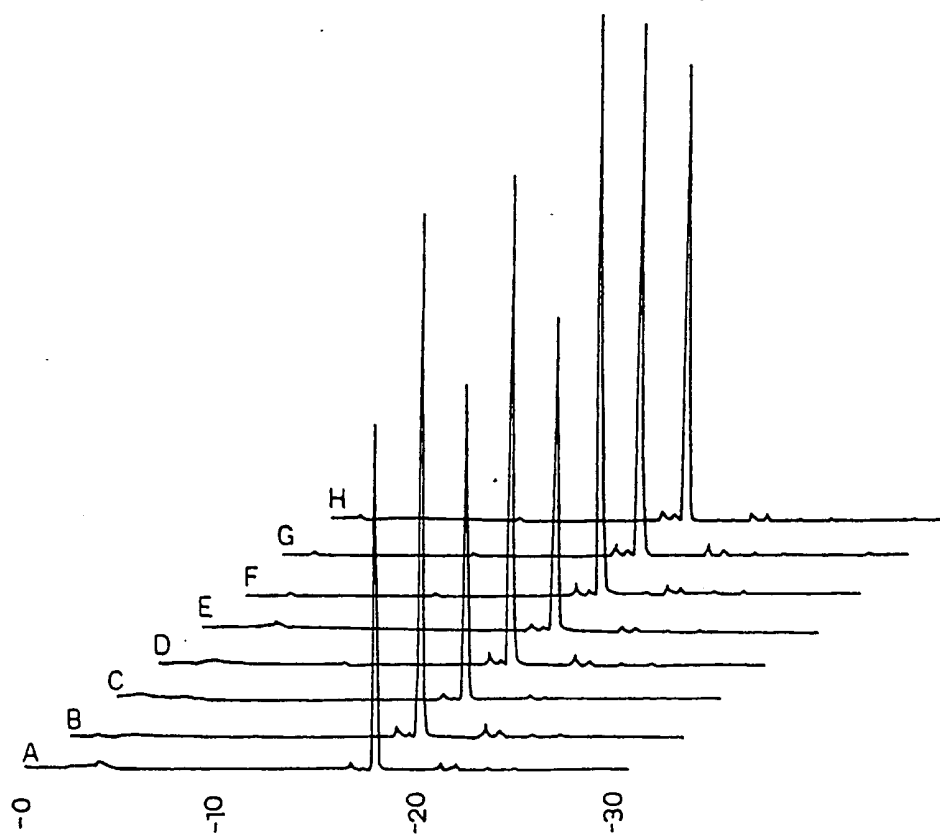


F I G . 1

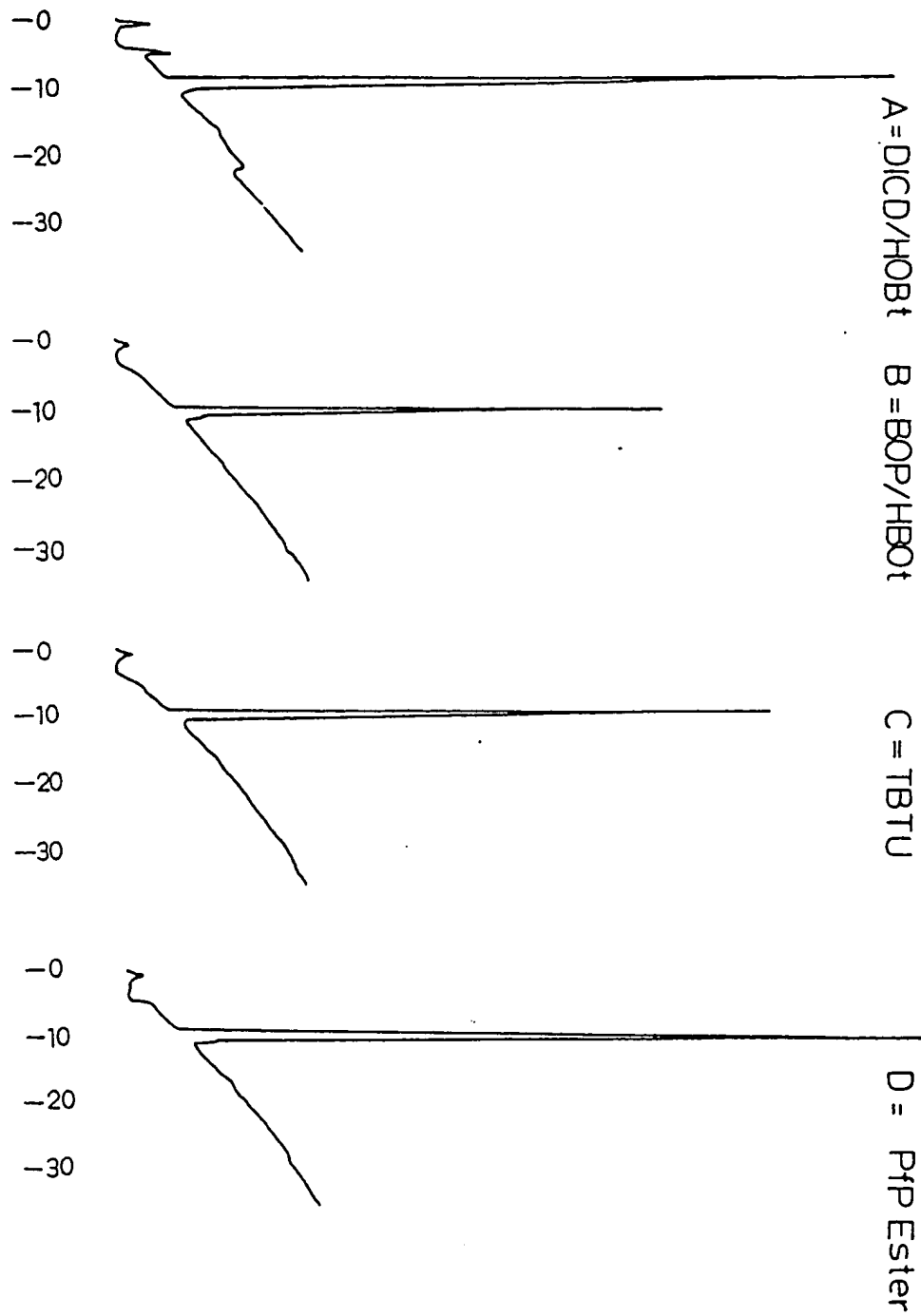


F I G . 2

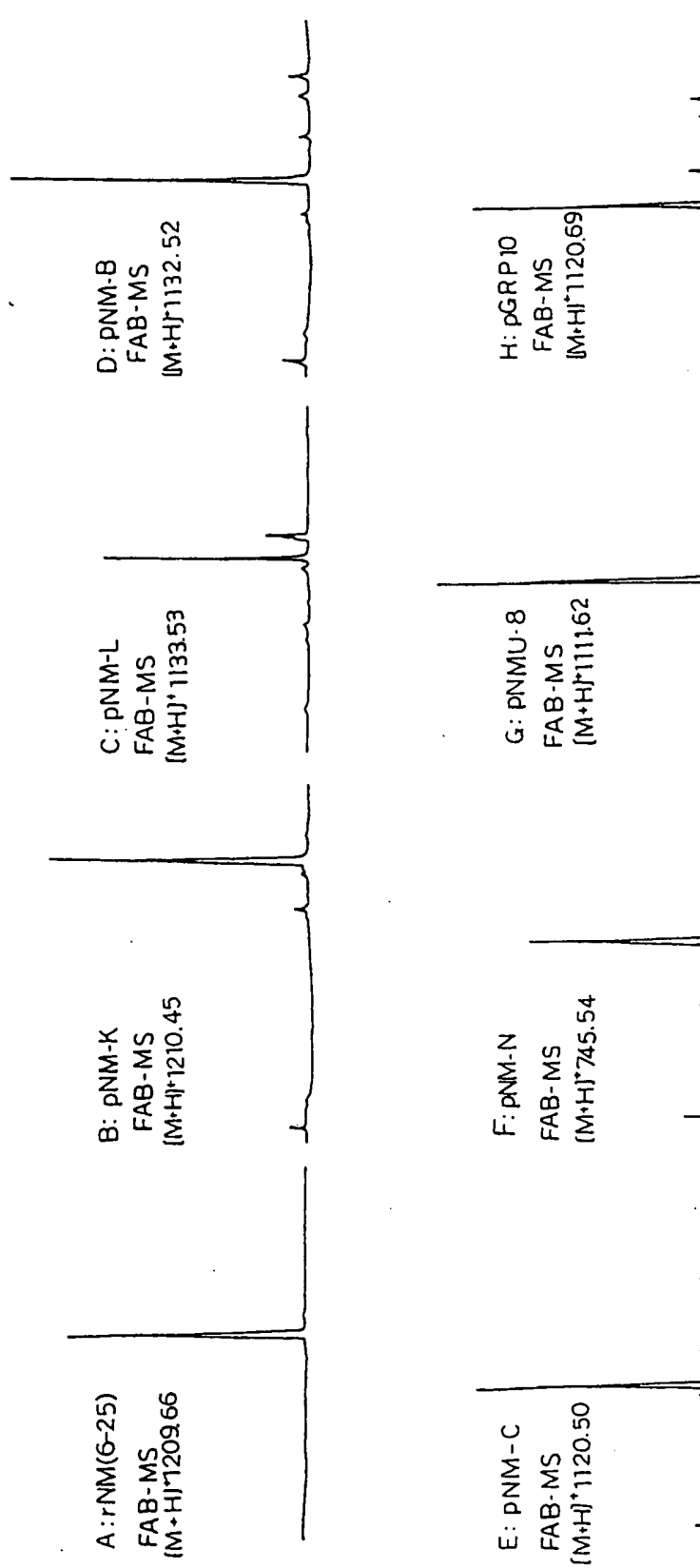




F I G . 4



F I G . 5



F I G . 6



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(54) **Apparatus for the simultaneous synthesis of different peptides.**

**EP 0 529 504 A3**

(57) The present invention is directed to the simultaneous multiple chemical synthesizer comprising a number of reaction vessels wherein each vessel has a filter in the bottom portion thereof, a number of needles wherein each needle is connected to a pair of an aspiration injection line of a reaction mixture and a gas supply line in charge of each reaction vessel and each needle does not touch resin, a number of arms which are horizontally and vertically movable and hold the respective needles, a pair of a bubbling gas line and a waste discharge line in charge of each reaction vessel, wherein each line is connected to the bottom portion of each reaction

vessel, a number of purge portions which move synchronously with said waste discharge lines, and means for washing the contact portions with the reaction reagents of the needles and the purge portions.



European Patent  
Office

# EUROPEAN SEARCH REPORT

Application Number  
EP 92 11 4165

DOCUMENTS CONSIDERED TO BE RELEVANT			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CL.5)
P, X	WO-A-91 13084 (BOEHRINGER INGELHEIM KG & GMBH) 5 September 1991 * the whole document *	1-4	B01J19/00 C07K1/04
A	EP-A-0 355 582 (BOEHRINGER INGELHEIM KG & GMBH) 28 February 1990 * the whole document *	1-4	
A	TETRAHEDRON, (INCL. TETRAHEDRON REPORTS) vol. 45, no. 24, 1989, OXFORD GB pages 7759 - 7764 G SCHNORRENBURG ET AL. 'fully automatic simultaneous multiple peptide synthesis in micromolar scale-rapid synthesis of series of peptides for screening in biological assays' * the whole document *	1-4	
A	DE-U-88 08 872 (H GAUSEPOHL) 29 December 1988 * the whole document *	3,4	
			TECHNICAL FIELDS SEARCHED (Int. CL.5)
			C07K B01J
The present search report has been drawn up for all claims			
Place of search THE HAGUE		Date of completion of the search 31 January 1994	Examiner Masturzo, P
<b>CATEGORY OF CITED DOCUMENTS</b> X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons * : member of the same patent family, corresponding document			

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